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(54) TIME: COMPOSITIONS AND METHODS FOR TREATING ATOPIC DERMATITIS, ANGIOEDEMA AND OTHER DISORDERS USING ANTHISTAMINES AND GLUCOCORTICOIDS

(57) Abstruct

Disclosed herein are compositions and methods for treating stopic dermatitis, angloedema, urdearia, altergio rhinitis and other such disorders. The compactions comprete therapeutically effective amounts of antihitsamines such as, for example, fortunations, and glucocardicodia such as, for example, beametaisons, for such freatment.

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COMPOSITIONS AND METHODS FOR TREATING ATOPIC DERMATITIS ANGIOEDEMA AND OTHER DISORDERS USING ANTHISTAMINES AND GLUCOCORTICOIDS

FIELD OF THE INVENTION

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The present invention generally relates to compositions and methods for treating atopic dermatitis, angioedema, urticaria, allergic rhinitis and other such disorders. It specifically discloses compositions comprising therapeutically effective amounts of antihistamines such as, for example, loratadine, and glucocorticoids such as, for example, betamethasone, for such treatment.

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BACKGROUND OF THE INVENTION

Atopic dermatitis is a chronic, itching, superficial inflammation of the skin, usually found in individuals with a history of allergic disorders. (The Merck Manual of Diagnosis and Therapy, D. Holvey ed., published by Merck & Co., Inc., Rahway, New Jersey, (1972) 1460). Angiocdema and urticaria are local wheals and erythema in the dermis and can be due to causes such as, for example, drug allergy, insect bites and the like, ibid, page 241. Atopic dermatitis is generally managed by applying ointments or pastes of topical corticosteroids. Itching is generally relieved by antihistamines, often in large doses. Initially useful medicaments may generally become ineffective and must be replaced. Acute urticaria is often managed by oral antihistamines; corticosteroid treatment may be occasionally necessary particularly when associated with angioedema. Topical corticosteroids are generally of no value.

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Products containing a combination of a steroid and an antihistamine are known and available. For example, a product containing chlorfeniramine maleate and paramethasone is available under the tradename DILARMINE® from Roche Pharmaceuticals, Nutley, New Jersey. However, the antihistamines in such products are typically sedating. There may be situations where sedating antihistamines are not acceptable.

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It would be desirable to find effective pharmaceutical compositions and methods of treatment for diseases such as atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergics, allergic contact dermatitis, seborrhoic dermatitis, neurodermatitis, allergic astiuna, ocular allergic manifestations such as conjunctivitis and iridociclitis, allergic reaction to insect stings and bites and other such disorders.

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It would be especially desirable to find compositions and methods of treatment for such diseases using an effective amount of a combination of one or more

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antihistamines, which are substantially non-sedating, with one or more glucocorticoids.

It would be additionally desirable to have such a combination composition and methods of treatment where the selected substantially non-sedating antihistamine(s) and glucocorticoid(s) are safe with low potential for systemic toxicity.

Other desires, objectives and advantages of the present invention will be apparent to those skilled in the art from the accompanying description and claims.

DESCRIPTION OF THE INVENTION

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The above-noted desires and objectives are addressed by the present invention which, in one embodiment, provides pharmaceutical compositions to treat diseases such as, for example, atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergies, allergic contact dermatitis, seborrhoic dermatitis, neurodermatitis, allergic asthma, coular allergic manifestations such as conjunctivitis and iridociclitis, allergic reaction to insect stings and bites and other such disorders. The composition comprises in combination: (i) a therapeutically effective amount of one or more substantially non-sedating antihistamine(s) and (ii) a therapeutically effective amount of one or more glucocorticoid or a suitable derivative thereof. The present invention additionally discloses a method for the treatment of the above-noted diseases in a mammalian organism in need of such treatment, such treatment comprising administering a pharmaceutical composition described above.

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The antihistamines useful in the practice of the present invention correspond to the general Formula I:

Formula I

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wherein X represents a halogen atom or a hydrogen atom; and Y represents hydrogen, -COOR, or -SO₂R₂, wherein R₁ represents a substituted or unsubstituted alkyl group, a substituted or unsubstituted develoalkyl group, a substituted or unsubstituted

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diacetate, desoxymetasone, desonide, cortivazol, corticosterone, cortisone, cloprednol, liethylaminoacetate), prednival, paramethasone, methylprednisolone, meprednisone, synthesized by the adrenal cortex of vertebrates and have anti-inflammatory activity. lumethasone, fludrocortisone, enoxolone, difluprednate, diflucortolone, diflorasone glucocorticoids and many are described in The Merck Index, Twelfth Ed., Merck & betamethasone, prednisolone, prednisone, flumethasone and hydrocortisone. Most cetoxypregnenolone, tralonide, diflurasone acetate, deacylcortivazol, budesonide, luocortolone, fluocortin butyl, fluocinonide, fluocinolone acetonide, flunisolode, deacylcortivazol oxetanone, and the like. Several of these compounds are known Many are well known. The glucocorticoids useful in the practice of the present Glucocorticoids generally belong to a class of steroid hormones that are clocortolone, clobetasone, clobetasol, chloroprednisone, cafestol, budesonide, lexamethasone, fluoromethalone, medrysone, triamcinolone; hydrocortisone, mazipredone, isoflupredone, halopredone acetate, halcinonide, formocortal, nvention include, for example, prednisolone, prednisone, betamethasone, prednicarbate, deflazacort, halomethasone, tixocortol, predinylidene (21l'urandrenolide, fluprednisolone, fluprednine acetate, fluperolone acetate, peclomethasone, ameinonide, allopreguane acetonide, alclometasone, 21-Do., Inc., Rahway, New Jersey (1996). Preferred glucocorticoids include preferred is betamethasone,

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antihistamine without clinically significant sedative effects. It has also been proven to be safe and effective for the treatment of several respiratory and dermatologic allergic considered to be a safe antihistamine as a H1 antagonist. It is also considered to be an Loratadine has been extensively studied for its antihistaminic effects and is hydrocortisone and has antiinflammatory properties. Betamethasone has been diseases. Betamethasone (Formula II) is a synthetic fluorinated derivative of

Formula II

Functions of Glucocorticoids in Stress and their Relation to Pharmacological Actions", loratadine and a glucocorticoid such as betamethasone has now been found to be a extensively used in clinic, as well as as a corticosteroid reference in several clinical highly effective medicament for treating allergic and related diseases stated above trials. A discussion can be found in, for example, A, Munck et al, "Physiological Endocr. Rev., Vol. 5 (1984), 25-44. A combination of an antihistamine such as without having a sedating effect.

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about 2-20 milligrams per dosage, preferably in about 2-10 milligrams and typically in pharmacologically acceptable salt or solvate is generally present in the composition in such as, for example, a pharmaceutically acceptable carrier. Still additional ingredients typically in about 0.03-0.5 milligram. Preferably the weight ratio of the glucocorticoid pharmaceutical compositions comprising a substantially non-sedating antihistamine about 3-7 milligrams. The glucocorticoid is generally present in the composition in and a glucocorticoid. Additionally, there may be other optional ingredients present may also be present, especially depending on the form of administration of the about 0.02-1 milligram per dosage, preferably in about 0.02-0.8 milligrams and As stated above, the present invention generally discloses novel pharmaceutical composition, as detailed later. The antihistamine or its and antihistamine is in the range between about 1:100 and 1:10.

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carrier is suitably selected with respect to the intended form of administration, i.e. oral carboxymethylcellulose, polyethylene glycol and waxes. Suitable lubricants that may diluents, excipients or carrier materials) may also be present in the composition. The benzoate, sodium acetate, sodium chloride, and the like. Disintegrants include starch, methylcellulose, guar gum and the like. Disinfectants include benzalkonium chloride ingredients such as, for example, sodium croscarmellose, may also be included where be mentioned for use in these dosage forms include, for example, boric acid, sodium charmaceutically acceptable inert carrier, such as lactose, starch, sucrose, cellulose, incorporated in the mixture. Suitable binders include starch, gelatin, natural sugars, magnesium stearate, dicalcium phosphate, calcium sulfate, mannitol, ethyl alcohol lablets, capsules (either solid-filled, semi-solid filled or liquid filled), powders for (liquid forms) and the like. Moreover, when desired or needed, suitable binders, administration in the form of tablets or capsules, the active ingredients (i.e., the antihistamine and the glucocorticoid) may be combined with any oral non-toxic constitution, oral gels, elixirs, solutions, syrups, suspensions, and the like, and and the like. Sweetening and flavoring agents and preservatives as well as other ubricants, disintegrating agents, disinfectants and coloring agents may also be As stated earlier, a pharmaceutically acceptable carrier (which includes consistent with conventional pharmaceutical practices. For example, for oral com sweeteners, natural and synthetic gums such as acacia, sodium alginate, appropriate

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In another embodiment, the present invention discloses a method of preparing a composition for use in the treatment of diseases such as, for example, atopic dernatitis, angioedema, uricaria, seasonal and allergic thinitis, food and drug allergies, allergic contact dermatitis, seborrhoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestatioms such as conjunctivitis and indociclitis, allergic reaction to insect strings and bites and other such disorders, with the composition comprising a therapeutically effective amount of one or more substantially non-sedating antihistamines, or a pharmaceutically acceptable salt or solvate of such antihistamine, and one or more therapeutically effective glucocorticoid, optionally in combination with a pharmaceutically acceptable carrier.

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In yet another embodiment, the present invention discloses a method of administering an effective treatment for diseases such as, for example, atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergies, allergic contact dermatitis, seborrhoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestations such as conjunctivitis and iridociclitis, allergic reaction to

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insect stings and bites and other such disorders, the administration comprising administering a pharmaceutical composition described above. The pharmaceutical compositions of the present invention can be administered depending upon the patient's age, sex, weight and severity of the condition being treated. Generally, the human oral dosage form containing the antihistamine and the carrier can be administered 1 or 2 times per day.

In a further embodiment, this invention discloses a method for the treatment of diseases such as, for example, atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergics, allergic contact dermatitis, sebornhoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestations such as conjunctivitis and irdociclitis, allergic reaction to insect stings and bites and other such disorders in a mammalian organism in need of such treatment, such treatment comprising administering a therapeutically effective amount of one or more substantially non-sedating antihistamines, or a pharmaceutically acceptable salt or solvate of such antihistamine, and one or more therapeutically effective glucocorticoid, optionally in combination with a pharmaceutically acceptable carrier.

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Additionally, the compositions of the present invention may be formulated in sustained release form to provide the rate controlled release of any one or more of the components or active ingredients to optimize the therapeutic effects. Suitable dosage forms for sustained release include layered tablets containing layers of varying disintegration rates or controlled release polymeric matrices impregnated with the active components and shaped in tablet form or capsules containing such impregnated or encapsulated porous polymeric matrices.

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Dosage form - refers to composition containing the antihistamine, the glucocorticoid and optionally a carrier formulated into a delivery system, i.e., tablet, capsule, oral gel, powder for constitution or suspension in association with inactive ingredients.

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Capsule - refers to a special container or enclosure made of methyl cellulose, polyvinyl alcohols, or denatured gelatins or starch for holding or containing compositions comprising the active ingredients and optionally the carrier. Hard shell capsules are typically made of blends of relatively high gel strength bone and pork skin gelatins. The capsule itself may contain small amounts of dyes, opaquing agents, plasticizers and preservatives.

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Tablet- refers to a compressed or molded solid dosage form containing the active ingredients with suitable diluents. The tablet can be prepared by compression

of mixtures or granulations obtained by wet granulation, dry granulation or by

Oral gels-refers to the active ingredients and the carrier dispersed or solubilized in a hydrophilic semi-solid matrix.

Powders for constitution refers to powder blends containing the active ingredients and the carrier and suitable diluents which can be suspended in water or inices.

Diluent - refers to substances that usually make up the major portion of the composition or dosage form. Suitable diluents include sugars such as lactose, sucrose, mannitol and sorbitol; starches derived from wheat, corn, rice and potato; and celluloses such as microcrystalline cellulose. The amount of diluent in the composition can range from about 2 to about 98% by weight of the total composition.

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Disintegrants - refers to materials added to the composition to help it break apart (disintegrate) and release the medicaments. Suitable disintegrants include starches; "cold water soluble" modified starches such as sodium carboxymethyl starch; natural and synthetic gums such as locust bean, karaya, guar, tragacanth and agar, cellulose derivatives such as methylcellulose and sodium carboxymethylcellulose; microcrystalline celluloses and cross-linked microcrystalline celluloses such as cross-microse such as againe acid and sodium alginate; clays such as bentonites; and effervescent mixtures. The amount of disintegrant in the composition can range from about 1 to about 15% by weight of the composition.

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Binders - refers to substances that bind or "glue" powders together and make them cohesive by forming granules, thus serving as the "adhesive" in the formulation. Binders add cohesive strength already available in the diluent or bulking agent. Suitable binders include sugars such as sucrose; starches derived from wheat, com, ricc and potato; natural gums such as acacia, gelatin and tragacantt; derivatives of seaweed such as alginio acid, sodium alginate and ammonium calcium alginate; cellulosic materials such as cellulose, methylcellulose, sodium carboxymethylcellulose and hydroxypropylmethylcellulose; pollyvinylpyrrolidone; and inorganics such as magnesium aluminum silicate. The amount of binder in the composition can range from about 2 to about 98% by weight of the composition.

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Lubricant - refers to a substance added to the dosage form to enable the tablet, granules, etc. after it has been compressed, to release from the mold or die by reducing friction or wear. Suitable lubricants include metallic stearates such as magnesium stearate, calcium stearate or potassium stearate, stearic acid; high melting point waxes; and water soluble lubricants such as sodium chloride, sodium benzoate, sodium

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acetate, sodium oleate, polyethylene glycols and d'I-leucine. Lubricants are usually added at the very last step before compression, since they must be present on the surfaces of the granules and in between them and the parts of the tablet press. The amount of lubricant in the composition can range from about 0.05 to about 5% by weight of the composition.

Glidents - materials that prevent caking and improve the flow characteristics of granulations, so that flow is smooth and uniform. Suitable glidents include silicon dioxide and talo. The amoint of glident in the composition can range from about 0.1% to about 5% by weight of the total composition.

Coloring agents - excipients that provide coloration to the composition or the dosage form. Such excipients can include food grade dyes and food grade dyes adsorbed onto a suitable adsorbent such as clay or aluminum oxide. The amount of the coloring agent can vary from about 0.1 to about 5% by weight of the composition.

Bioavailability - refers to the rate and extent to which the active drug ingredient or therapeutic moiety is absorbed into the systemic circulation from an administered dosage form as compared to a standard or control, as well as to topical bioavailability. Conventional methods for preparing tablets are known. Such methods include

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Convenieura incurous for preparing tables are known. Such intendes include dry methods such as direct compression and compression of granulation produced by compaction, or wet methods or other special procedures.

The phrase "therapeutically effective amount" means that amount of the active ingredients which provides a therapeutical benefit in the treatment or management of the diseases stated above by the present inventive composition.

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The magnitude of a prophylactic or therapeutic dose of the active ingredients in the acute or chronic management of the targeted disease or condition will vary with the severity of the condition to be treated and the route of administration. The dose, and perhaps the dose frequency, will also vary according the age, body weight, and response of the individual patient. Suitable total daily dose ranges can be readily determined by those skilled in the art. The dose may be administered in single or divided doses orally, topically, transdermally, or locally by inhalation.

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It is further recommended that children, patients aged over 65 years, and those with impaired renal or baptic function initially receive low doses, and that they then be titrated based on individual response(s) or blood level(s). Further, it is noted that the clinician or treating physician will know how and when to adjust, interrupt, or terminate therapy in conjunction with individual patient response.

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Any suitable route of administration may be employed for providing the patient with an effective dosage of the active ingredients according to the methods of

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the present invention. Some such routes are, for example, oral, intraoral, rectal, parenteral, epiculaneous, transdermal, subcutaneous, intramuscular, intranasal, sublingual, intradural, intraocular, intrarespiratory, oral or nasal inhalation and the like. Oral administration is preferred.

The term "pharmaccutically acceptable salt" refers to a salt prepared from pharmaccutically acceptable non-toxic acids or bases including inorganic acids or bases or organic acids or bases. Examples of such inorganic acids are hydrochloric, hydrodic, sulfuric, and phosphoric. Appropriate organic acids may be selected, for example, from aliphatic, aromatic, carboxylic.and sulfonic classes of organic acids, examples of which are formic, acetic, propionic, succinic, glycolic, glucuronic, maleic, furoic, glutamic, benzoic, anthranilic, salicylic, phenylacetic, mandelic, embonic (pamoic), methanesulfonic, ethanesulfonic, pantothenic, benzenesulfonic, stearic, sulfanilic, algenic, and galacturonic. Examples of such inorganic bases include metallic salts made from aluminum, calcium, lithium, magnesium, potassium, sodium, and zinc. Appropriate organic bases may be selected, for example, from N,N-dibenzylethylenediamine, chloroprocaine, choline, diethanolamine, ethylenediamine, meglumaine (N-methylgulcaine), lysine and procaine. A similar meaning is given to the term "pharmaceutically acceptable solvate" which, however, includes a solvent, water and the ikke as the solvating medium.

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As stated before, dosage forms include tablets, troches, dispersions, suspensions, subtitions, capsules, patches, syrups, elixirs, gels, powders, magmas, lozenges, ointments, creams, pastes, plasters, lotions, discs, suppositories, nasal or oral sprays, aerosols and the like. Because of their ease of administration, tablets and capsules represent the most advantageous oral dosage unit form, in which case solid pharmaceutical carriers are employed. If desirable, tablets may be coated by standard aqueous or nonaqueous techniques. Another preferred dosage form is as liquid or solution, comprising the active ingredients along with any additional optional ingredient or ingredients in a pharmaceutically acceptable carrier which is preferably a liquid.

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Pharmaceutical compositions for use in the methods of the present invention may be prepared by any of the methods of pharmacy, but all methods include the step or steps of bringing into association the active ingredients and any optional ingredient or ingredients, carrier and the like. Generally stated, the compositions are prepared by uniformly and intimately admixing the active ingredients with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product into the desired presentation.

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For example, a tablet may be prepared by compression or molding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing in a suitable machine the active ingredients in a free-flowing form such as powder or granules, optionally mixed with a binder, lubricant, inert diluent, surface active or dispersing agent. Molded tablets may be made by molding, in a suitable machine, a mixture of the powdered compound moistened with an inert liquid diluent.

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An example composition containing loratadine and betamethasone for a tablet and preparation of a tablet by a compression molding process may be illustrated as in the following Table 1:

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Name of the Ingredient	Concentration	Concentration range (milligram/tablet)
Betamethasone		0.1-0.5
Loratadine		2-10
Lactose monohydrate (LACTOSE FASTFLO®)	FASTFLO®)	55-290
Sodium Croscarmellose		.0.8-4
Magnesium stearate		0.4-1
Tablet weight:		60-300 tilg.
The above-stated ingredients may be admixed in any suitable order and converted into	ked in any suitat	ile order and converted into
a tablet by suitable methods such as, for example, the methods stated earlier. Thus, in	cample, the meth	ods stated earlier. Thus, in
one example, betamethasone (0.25 mg) and loratadine (5 mg) were premixed with one	l loratadine (5 m	g) were premixed with one
portion (70 mg) of Lactose FASTFLO® (available from Foremost Farm USA,	vailable from Fo	remost Farm USA,
Baraboo, Wisconsin). The mix was then passed through a Quadro CoMil® mill (a sieve	ussed through a	Quadro CoMiil® mill (a sieve
mill available from Quadro, Waterloo, Ontario, Canada) equipped with a 20 mesh	ario, Canada) eq	uipped with a 20 mesh
screen. The remaining portion of lactose FASTFLO® (71.75 mg) and croscarmellose	ASTFLO® (71.7;	5 mg) and croscarmellose
sodium (2 mg) were then added and blended. Magnesium stearate (1 mg) was then	d. Magnesium s	tearate (1 mg) was then
mixed in and blended well. The mixture was then compressed in a rotary tablet press	as then compress	sed in a rotary tablet press
to make tablets. A tablet weighing about 150 milligrams may be prepared with the	50 milligrams m	ay be prepared with the
above-noted composition.		

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Alternatively, a tablet may be prepared by a wet granulation method. An example composition containing loratadine and betamethasone for a tablet and preparation of a tablet by a wet granulation process may be illustrated as in the following Table II. In Table II, AVICEL PH301° is a microcoystalline cellulose, available from FMC Corporation Pharmaceutical Division, City of Industry, California; Kollidon VA64° is a modified polyvinyl pyrrolidone available from BASF Mexicana S.A de C.V. Av, de los Deportes C.P., Mexico D.F.; Kollidon CL° is

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another modified polyvinyl pyrrolidone also available from BASF Mexicana S.A de C.V. Av, de los Deportes C.P., Mexico D.F.

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Table II

Name of the Ingredient	Concentration range (milligram/tablet)
Betamethasone (micronized)	0.1-0.5
Loratadine (micronized)	2-10
Cellulose microcrystalline (e.g., AVICEL PH301°) 55-290	VICEL PH301°) 55-290
Kollidon VA64®	1-5
Kollidon CL®	2-10
Magnesium stearate	0.1-1
Ethyl alcohol	as needed (about 20-40 weight % based
	on total formulation)

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Tablet weight:

In the illustrative process, in a suitable container, sufficient alcohol was taken to dissolve loratadine (5 mg), betamethasone (0.25 mg) and Kollidon VA64® (3 mg) and the three ingredients were dissolved in it. In a separate mixer, cellulose (84.65 mg) and Kollidon CL® (7 mg) were mixed for about 15 minutes and this mix was then blended and granulated with the alcohol solution and mixed until a uniform granulate was formed. The granulate was then spreaded on trays and dried in an oven until a 1.5-2.5% moisture level was obtained. The granulate was then passed through a 25 mesh screen, charged into a mixer and mixed well with magnesium stearate (0.1 mg) for about 2 minutes. This mix was then compressed into tablets using a tablet press machine and 4 inch deep concave round punches.

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Various modifications of the invention in addition to those shown and described herein will be apparent to those skilled in the art from the foregoing description. Such modifications are intended to fall within the scope of the invention and appended claims. Furthermore, even though the above-stated illustrative examples are prepared using loratadine and betamethasone as the active ingredients, it should not be considered as limiting the scope of the invention in any way; other suitable substitution of the active ingredients may be made using the earlier stated lists of suitable compounds. Modifications of the active ingredients and the other ingredients as well as the process suitably are also to be considered as falling within the scope of the invention, description and appended claims.

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CLAIMS

What is claimed is:

- 1. A pharmaceutical composition for treating atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergies, allergic contact dermatitis, seborthoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestations such as conjunctivitis and iridocicilitis, and allergic reaction to insect stings and bites, said composition comprising: (i) a therapeutically effective amount of one or more substantially non-sedating antihistamines or a pharmaceutically acceptable salt or solvate thereof, and (ii) a therapeutically effective amount of one or more glucocorticoid.
- The composition of claim 1 for treating atopic dermatitis.

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- The composition of claim 1 for treating angioedema.
- The composition of claim 1, wherein said antihistamine corresponds to the general formula:

wherein X represents a halogen atom or a hydrogen atom; and Y represents hydrogen, -COOR₁ or -SO₂R₂, wherein R₁ represents a substituted or unsubstituted alkyl group, a substituted or unsubstituted or unsubstituted alkyl group, or a substituted or unsubstituted alkenyl group, or a substituted or unsubstituted or unsubstituted anyl group, or a substituted or unsubstituted cycloalkyl group, or substituted or unsubstituted anyl group as well as optical isomers and mixtures of said antihistamine, with said substituents being a C1-C6 alkyl, aralkyl, alkylaryl, aryl and cycloalkyl.

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5. The composition of claim 4, wherein X is a halogen and Y is hydrogen or COOR₁₁, wherein R₁ is the same as in claim 4.

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- The composition of claim 5, wherein X is Cl and Y is -COOC₂H₅, said antihistamine being known as loratadine.
- The composition of claim 5, wherein X is Cl and Y is hydrogen, said antihistamine being known as desloratadine.

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8. The composition of claim 1, wherein said glucocorticoid is selected from the group consisting of prednisolone, prednisone, betamethasone, dexamethasone, fluoromethalone, medrysone, triamcinolone, hydrocortisone, prednicarbate, deflazacort, halomethasone, tixocotol, predinylidene (21-diethylaminoacetate), prednival, paramethasone, methylprednisolone, meprednisone, mazipredone, isoflupredone, halopredone acetate, fluoromode, formocortal, flurandrenolide, fluprednisolone, fluoromodone acetate, fluoromode, fluoromodine, fluoromodine acetate, fluoromodole, fluoromodine, fluoromodine, fluoromodine, fluoromodine, coprednol, clocortolone, fluorocortisone, enoxolone, difluprednate, diflucortolone, cloprednol, clocortolone, clobetasone, desonide, cortivazol, corticosterone, cortisone, cloprednol, clocortolone, clobetasone, clobetasol, chloroprednisone, cafestol, budesonide, beclomethasone, ancinonide, allopregnane acetonide, alchometasone, 21-acetoxypregnenolone, tralonide, diflurasone acetate, deacylcortivazol, budesonide, deacylcortivazol oxetanone, and mixtures thereof.

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The composition of claim 8, wherein said glucocorticoid is betamethasone.
 The composition of claim 1, additionally containing one or more ingredients selected from the zrow consisting of a pharmaceutically accentable carrier. binder.

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10. Its composition of claim 1, additionally containing one or more ingredients selected from the group consisting of a pharmaceutically acceptable carrier, binder, lubricant, disintegrating agent, disinfectant, coloring agents, flavoring agent and preservative.

 The composition of claim 10, wherein said additional ingredient is a pharmaceutically acceptable carrier.

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12. The composition of claim 11, wherein said pharmaceutically acceptable carrier is selected from the group consisting of lactose, sucrose, sugar, cellulose, magnesium stearate, dicalcium phosphate, calcium sulfate, mannitol, ethyl alcohol and mixtures thereof.

13. The composition of claim 12, wherein said carrier is lactose.

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14. The composition of claim 12, wherein said carrier is magnesium stearate.

15. The composition of claim 1, additionally comprising croscarmellose sodium.

The composition of claim 1, being present in the form of a tablet.

The composition of claim 1, being present in the form of a capsule.

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The composition of claim 1, being present in the form of a liquid.
 The composition of claim 18, wherein said liquid is a solution of said

 The composition of claim 18, wherein said liquid is a solution of said composition in a suitable solvent. The composition of claim 16, wherein said tablet is prepared by compression.

21. The composition of claim 16, wherein said tablet is prepared by granulation.

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22. The composition of claim 1, wherein said antihistamine is present in amounts in the range 2-20 milligrams per dosage. 23. The composition of claim 1, wherein said antihistamine is present in amounts in the range 2-10 milligrams per dosage.

The composition of claim 1, wherein said antihistamine is present in about 3-7
milligrams amounts per dosage.

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 The composition of claim 1, wherein said glucocorticoid is present is about 0.02-1 milligram per dosage. The composition of claim 1, wherein said glucocorticoid is present is about 0.02-0.8 milligram per dosage.

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 The composition of claim 1, wherein said glucocorticoid is present is about 0.03-0.5 milligram per dosage. 28. The composition of claim 1, wherein said glucocorticoid and said antihistamine are present in a respective weight ratio range between 1:100 and 1:10.

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29. A method for the treatment of atopic dermatitis, angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergics, allergic contact dermatitis, seborthoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestations such as conjunctivitis and iridociclitis, and allergic reaction to insect stings and bites, in a mammalian organism in need of such treatment, said method comprising administering to said organism a composition comprising; (i) a therapeutically effective

amount of one or more substantially non-sedating antihistantines or a pharmaceutically acceptable salt or solvate thereof, and (ii) a therapeutically effective amount of one or more glucocorticoid.

30. The method of claim 29, wherein said antihistamine corresponds to the general

wherein X represents a halogen atom or a hydrogen atom; and Y represents hydrogen, $-\text{COOR}_1$ or $-\text{SO}_2R_2$, wherein R₁ represents a substituted or unsubstituted alkyl group,

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cycloalkyl group, or substituted or unsubstituted aryl group as well as optical isomers unsubstituted heterocyclic ring; and R2 represents a substituted or unsubstituted a substituted or unsubstituted cycloalkyl group, a substituted or unsubstituted alkenyl group, a substituted or unsubstituted aryl group, or a substituted or and mixtures of said antihistamine.

The method of claim 30, wherein X is Cl and Y is -COOC2H5, said antihistamine being known as loratadine.

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The method of claim 29, wherein said glucocorticoid is selected from the group outyl, fluocinonide, fluocinolone acetonide, flunisolode, flumethasone, fludrocortisone, ulopregnane acetonide, alciometasone, 21-acetoxypregnenolone, tralonide, diflurasone lesonide, cortivazol, corticosterone, cortisone, cloprednol, clocortolone, clobetasone, supreduisolone, flupreduine acetate, fluperolone acetate, fluocortolone, fluocortin slobetasol, chloroprednisone, cafestol, budesonide, beclomethasone, ameinonide, anoxolone, difluprednate, diflucortolone, diflorasone diacetate, desoxymetasone, cetate, deacylcortivazol, budesonide, deacylcortivazol oxetanone, and mixtures deflazacort, halomethasone, tixocortol, predinylidene (21-diethylaminoacetate), prednival, paramethasone, methylprednisolone, meprednisone, mazipredone, soflupredone, halopredone acetate, halcinonide, formocortal, flurandrenolide, fluoromethalone, medrysone, triamcinolone, hydrocortisone, prednicarbate, consisting of prednisolone, prednisone, betamethasone, dexamethasone, hereof.

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The method of claim 32, wherein said glucocorticoid is betamethasone.

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allergic manifestations such as conjunctivitis and iridociclitis, and allergic reaction to insect stings and bites, said composition comprising: (i) a therapeutically effective unount of loratadine or a pharmaceutically acceptable salt or solvate thereof, (ii) a angioedema, urticaria, seasonal and allergic rhinitis, food and drug allergies, allergic contact dermatitis, seborrhoic dermatitis, neurodermatitis, allergic asthma, ocular A pharmaceutical composition for the treatment of atopic dermatitis, herapeutically effective amount of betamethasone; and (iii) lactose.

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The composition of claim 34, additionally comprising magnesium stearate and croscarmellose sodium.

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such as conjunctivitis and iridociclitis, and allergic reaction to insect stings and bites, in seborrhoic dermatitis, neurodermatitis, allergic asthma, ocular allergic manifestations seasonal and allergic rhinitis, food and drug allergies, allergic contact dermatitis, A method for the treatment of atopic dermatitis, angioedema, urticaria, mammalian organism in need of such treatment, said method comprising

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administering to said organism a composition comprising: (i) a therapeutically effective amount of loratadine or a pharmaceutically acceptable salt or solvate thereof; (ii) a therapeutically effective amount of betamethasone; and (iii) lactose.

The method of claim 36, wherein said composition additionally comprises magnesium stearate and croscarmellose sodium. 37.

INTERNATIONAL SEARCH REPORT

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# GAMBLE) ines 15–30, ines 15		704 206 A (REGENOLD JU -11 1996 (1996-04-03) claims 1 and 29*	ERGEN OR)		1-37
	······································	780 127 A (PROCTER & G une 1997 (1997-06-25) abstract, page 3, line cample 1, claim 1*	AMBLE) s 15-30, page		1-37
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INTERNATIONAL SEARCH REPORT

PC., US 99/04502 CiContinuation OOCUMENTS CONSIDERED TO BE RELEVANT
Chalon of document, with indication, when appropriate, of the relevant passages
JEAL W. ET AL.: "Triamcinolone acetonide. A review of its pharmacological properties and therapeutic efficacy in the management of allergic rinnits." DRUGS, vol. 55, no. 2, February 1997 (1997-02), vol. 55, no. 2, February 1997 (1997-02), cf. pages 257-280, XP002121955 cf. page 258, lines 16-21, page 260, 19-21, page 273, 4th para, on the 19ftt-handed col., page 274, left col.,
RATMER F. H., ET AL.: "A comparison of the efficacy of fluticasone propionate aqueous masal spray and loratadine, alone and in combination, for the treatment of seasonal allergic rhinitis." JOHNAL OF FAMILY PRACTICE, VOUNAL OF FAMILY PRACTICE, VOUNAL OF FAMILY PRACTICE, Pages 118-125, XP002121956 Coi., Dage 118, "background", right-handed coi., Background", right-handed coi., 1st para., page 120, table 1, page 121, figure 1 and page 124, left-hand col., 3rd para.,*

page 2 of 2

INTERNATIONAL SEARCH REPORT

Patent family member(s) 2201358 A 9610389 A 19536244 A 19536245 A		fon	formation on patent family members	mbers	PC., US 99/04502	PC., US 99/04502
A 03-04-1996 CA 2201358 A WO 9610389 A DE 19536244 A DE 19536245 A DE 19536245 A	Patent document cited in search report		Publication date	u	atent family member(s)	Publication date
19536246 A 5958379 A	EP 0704206	A	03-04-1996	S S S S S S S S S S S S S S S S S S S	2201358 A 9610389 A 19536244 A 19536245 A 19536246 A 5958379 A	11-04-1996 11-04-1996 04-04-1996 04-04-1996 28-09-1999
EP 0780127 A 25-06-1997 NONE	EP 0780127	A	25-06-1997	NONE		

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